Amendments to the Claims

This listing of claims will replace all prior versions and listing of claims in the application.

- 1-28. (Canceled)
- 29. (Currently amended) A pharmaceutical formulation comprising a core <u>containing</u>, as <u>ingredients</u>, and core materials, wherein the core materials comprise therapeutically effective amounts of an IBAT inhibitor compound, and a bile acid binder and optional pharmaceutically acceptable excipients, and wherein the core <u>ingredient</u> material comprising the bile acid binder is coated with a layer for the targeted release of the bile acid binder in the colon.
- 30-32. (Canceled)
- 33. (Currently amended) A pharmaceutical formulation comprising a core <u>containing</u>, as <u>ingredients</u>, and core materials, wherein the core materials comprise therapeutically effective amounts of an IBAT inhibitor compound, and a bile acid binder and optional pharmaceutically acceptable excipients, and wherein the core <u>ingredient material</u> comprising the IBAT inhibitor compound is coated with a layer for the targeted release of the IBAT inhibitor compound in the ileum and the core <u>ingredient material</u> comprising <u>the</u> bile acid binder is coated with a layer for the targeted release of the bile acid binder in the colon.
- 34. (Previously presented) A method for the therapeutic treatment of a subject suffering from, or susceptible to hypercholesterolemia, wherein the method comprises administering to the subject a therapeutically effective amount of an IBAT inhibitor compound and a bile acid binder, wherein the bile acid binder is administered for the therapeutic treatment of diarrhea during administration of the IBAT inhibitor.
- 35. (Currently amended) A method for the therapeutic treatment of a subject suffering from, or susceptible to, diarrhea during administration of an IBAT inhibitor compound, comprising administering to the subject a pharmaceutical formulation comprising a core containing, as ingredients, and core materials, wherein the core materials comprise therapeutically effective amounts of a bile acid binder and optional pharmaceutically

- acceptable excipients, and wherein the core ingredient material comprising the bile acid binder is coated with a layer for targeted release of the bile acid binder in the colon.
- 36. (Previously presented) The pharmaceutical formulation according to claim 29 or 33, wherein the IBAT inhibitor compound is a low permeability drug as defined in the FDA Biopharmaceutical Classification System.
- 37. (Previously presented) The pharmaceutical formulation according to claim 29 or 33, wherein the bile acid binder is a resin.
- 38. (Previously presented) The pharmaceutical formulation according to claim 29 or 33, wherein the IBAT inhibitor compound and the bile acid binder are administered simultaneously, separately or sequentially.
- 39. (Previously presented) The pharmaceutical formulation according to claim 29 or 33, wherein the IBAT inhibitor compound comprises a benzothiazepine having IBAT inhibiting properties.
- 40. (Previously presented) The pharmaceutical formulation according to claim 39, wherein the benzothiazepine is 1,4-benzothiazepine or a 1,5-benzothiazepine.
- 41. (Previously presented) The method according to claim 34 or 35, wherein the IBAT inhibitor compound is a low permeability drug as defined in the FDA Biopharmaceutical Classification System.
- 42. (Previously presented) The method according to claim 34 or 35, wherein the bile acid binder is a resin.
- 43. (Previously presented) The method according to claim 34 or 35, wherein the IBAT inhibitor compound and the bile acid binder are administered simultaneously, separately or sequentially.
- 44. (Previously presented) The method according to claim 34 or 35, wherein the IBAT inhibitor compound comprises a benzothiazepine having IBAT inhibiting properties.
- 45. (Previously presented) The method according to claim 44, wherein the benzothiazepine is 1,4-benzothiazepine or a 1,5-benzothiazepine.